

# MojoSort™ Mouse Neutrophil Isolation Kit Column Protocol

DOI

dx.doi.org/10.17504/protocols.io.7bjhikn



### Sam Li<sup>1</sup>

<sup>1</sup>BioLegend

BioLegend

Tech. support email: tech@biolegend.com



Sam Li

BioLegend

### Create & collaborate more with a free account

Edit and publish protocols, collaborate in communities, share insights through comments, and track progress with run records.

Create free account



DOI: https://dx.doi.org/10.17504/protocols.io.7bjhikn

External link: https://www.biolegend.com/protocols/mojosort-mouse-neutrophil-isolation-kit-column-protocol/4770/

Protocol Citation: Sam Li . MojoSort™ Mouse Neutrophil Isolation Kit Column Protocol. protocols.io <a href="https://dx.doi.org/10.17504/protocols.io.7bjhikn">https://dx.doi.org/10.17504/protocols.io.7bjhikn</a>

**License:** This is an open access protocol distributed under the terms of the **Creative Commons Attribution License**, which permits unrestricted use, distribution, and reproduction in any medium, provided the original author and source are credited

Created: September 13, 2019



Last Modified: September 13, 2019

Protocol Integer ID: 27723

**Keywords:** MojoSort<sup>™</sup>, nanobeads, column, cell separation, less antibody cocktail that with other commercial supplier, less antibody cocktail, antibody cocktail, best dilution factor, dilution, antibody, streptavidin nanobead, fewer bead, nanobead, used separation column

#### Abstract

BioLegend MojoSort<sup>™</sup> nanobeads work in commonly used separation columns, based on our internal research as well as validation by external testing by academic labs. This simple protocol consists of following the MojoSort<sup>™</sup> protocol to label the cells with **pre-diluted** MojoSort<sup>™</sup> reagents and using the columns as indicated by the manufacturer.

**Note:** Due to the properties of our beads, it may be possible to use far fewer beads and less antibody cocktail that with other commercial suppliers. We recommend a titration to find the best dilution factor. However, as a general rule, dilutions ranging from 1:2 to 1:10 for the antibody cocktail can be used. Dilutions ranging from 1:5 to 1:20 for the Streptavidin Nanobeads can be used. Please contact BioLegend Technical Service (tech@biolegend.com) if further assistance is needed.

#### Guidelines

MojoSort™ magnetic particles can be used with other commercially available magnetic separators, both free standing magnets and column-based systems. Because MojoSort™ protocols are optimized for the MojoSort™ separator, the protocols may need to be adjusted for other systems. Please contact BioLegend Technical Service (tech@biolegend.com) for more information and guidance. We do not recommend using MojoSort™ particles for BD's IMag™ or Life Technologies' DynaMag™.

#### **Materials**

**MATERIALS** 

**⊠** MojoSort<sup>™</sup> Buffer **BioLegend Catalog** #480017

Additional reagents:

- -commercially available cell separation columns
- -5 mL polypropylene tubes
- -70 μm cell strainer

This protocol works with the following MojoSort™ Kits (cat#):

480057, 480058

# **Troubleshooting**



- 1 Prepare cells from your tissue of interest or blood without lysing erythrocytes.
- In the final wash of your sample preparation, resuspend the cells in MojoSort™ Buffer by adding up to 4 mL in a 5 mL (12 × 75 mm) polypropylene tube.

  Note: Keep MojoSort™ Buffer on ice throughout the procedure.
- 3 Filter the cells with a 70 µm cell strainer, centrifuge at 300xg for 5 minutes, and resuspend in an appropriate volume of MojoSort™ Buffer. Count and adjust the cell concentration to 1 × 10<sup>8</sup> cells/mL by adding MojoSort™ Buffer.

5m

Aliquot 100 μL (10<sup>7</sup> cells) into a new tube. Add **10 μL of the pre-diluted Biotin-Antibody Cocktail**. Mix well and **incubate on ice for 15 minutes**. Scale up the volume if separating more cells. For example, add 100 μL of pre-diluted Antibody Cocktail for separating 1 × 10<sup>8</sup> cells in 1 ml of MojoSort™ Buffer. When working with less than 10<sup>7</sup> cells, use indicated volumes for 10<sup>7</sup> cells.

15m

Wash the cells by adding MojoSort™ Buffer up to 4 mL. Centrifuge the cells at 300xg for 5 minutes.

5m

6 Discard the supernatant and resuspend cells in 100 µL of MojoSort™ Buffer.

15m

Vortex the Streptavidin conjugated Nanobeads (to resuspend) at max speed, 5 touches, and prepare the dilutions to test. Add **10 μL of pre-diluted Streptavidin Nanobeads**. Mix well and **incubate on ice for 15 minutes**. Scale up the volume accordingly if separating more cells. For example, add 100 μL of pre-diluted Nanobeads for separating 1 × 10<sup>8</sup> cells in 1 ml of MojoSort™ Buffer. When working with less than 10<sup>7</sup> cells, use indicated volumes for 10<sup>7</sup> cells.

5m

- 8 Wash the cells by adding MojoSort™ Buffer up to 4 mL. Centrifuge the cells at 300xg for 5 minutes.
- 9 Discard the supernatant.
- 10 Resuspend the cells in appropriate amount of MojoSort™ Buffer and proceed to separation. At least 500 µL is needed for column separation.

**Note:** There are several types of commercially available columns, depending on your application. Choose the one that fits best your experiment:



	Max. number of labeled cells	Max. number of total cells	Cell suspension volume	Column rinse volume	Cell wash volume	Elution volume
Small Capacity	1 x 10 <sup>7</sup>	2 x 10 <sup>8</sup>	500µL for up to 10 <sup>8</sup> cells	1ml	1 ml	1 ml
Medium Capacity	1 x 10 <sup>8</sup>	2 x 10 <sup>9</sup>	500µL for up to 10 <sup>9</sup> cells	3ml	3 ml	5 ml
Large Capacity	1 x 10 <sup>9</sup>	2 x 10 <sup>10</sup>	500µL for up to 10 <sup>10</sup> cells	20-50ml	30 ml	20 ml

# Example of magnetic separation with medium capacity columns:

- 11 Place the column in a magnetic separator that fits the column.
- 12 Rinse the column with 3 mL of cell separation buffer.
- 13 Add the labeled cell suspension in at least 500 µL of buffer to the column through a 30 µm filter and collect the fraction containing the unlabeled cells. These are the cells of interest; do not discard.
- 14 Wash the cells in the column **1 time** with 3 mL of buffer and collect the fraction containing the untouched cells. Combine with the collected fraction from step 3.
- 15 If desired, the labeled cells can be collected by taking away the column from the magnet and place it on a tube. Then add 5 mL of buffer and flush out the magnetically labeled fraction with a plunger or supplied device. The labeled cells may be useful as staining controls, to monitor purity/yield, or other purposes.